Reinforcement Learning: An Introduction

Summary Notes

Mrinank Sharma

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1 Introduction

Other than the **agent** and the **environment**, there are four main subelements of an RL system:

- Elements of Reinforcement Learning
- 1. The *policy* maps perceived states of the environment to actions. This could be a lookup table or a search process.
- The reward signal defines the goal of a reinforement learning problem. The environment sends rewards to an agent that has the sole objective of maximising this reward.
- 3. The *value function* specifies what is good in the long run. The *value* of a state is the total amount of reward an agent can expect to accumulate over teh future starting from that state.
- 4. Some RL systems have *models* of the environment that can mimic the behaviour of the environment. Models are used for planning.

Whilst rewards determine the immediate, intrisic desirability of environmental states, values indicate the long-term desirability of states after taking into account which states are likely to follow. We choose actions based on values, even though values are derived from rewards.

It is possible to learn policies without estimating value functions by using evolutionary methods. However, they tend to ignore much of the useful structure and are not well suited for RL tasks as they ignore relevant information, such as the states which are passed and the actions which are selected.

Rewards and Value Functions

Evolutionary Methods

2 Multi-armed Bandits

The most important feature distinguishing reinforcement learning from other types of learning is that it uses training information that **evaluates** the actions taken rather than **instructs** by giving correct actions.

This form of feedback creates the need for active exploration.

Definition 2.1 (k-armed Bandit) In the k-armed bandit problem, you are repeatedly faced with a choice among k different actions. After each choice, a numerical reward is received from a **stationary** probability distribution associated with the selected action. The objective is to maximise the expected total reward over some time period.

k-armed bandits

The value of action a is the expected reward given a was chosen:

$$q_*(a) = \mathbb{E}[R_t | A_t = a]. \tag{1}$$

If we knew the values, problem solved - just pick the action with the maximum value. However, we do not know these. Denote the *estimated* value as $Q_t(a)$. A simple way to estimate the action-value is a *sample average*, which can be calculated incrementally.

Note that purely greedy methods are best if there is no noise in the reward signal. With ϵ -greedy algorithms, the best value of ϵ will depend on the amount of noise, which

Reward Variance

determines the number of samples for the sample average to converge. However, if the problem is *non-stationary*, we still need to explore.

If the bandit is non-stationary, we might use the following rule for value estimates:

Non-stationary Problems

$$Q_{n+1}(a) = Q_n(a) + \alpha_n(a) \left[R_n - Q_n(a) \right]. \tag{2}$$

It is known that convergence with probability 1 happens if the following conditions hold:

Convergence Guarantees

$$\sum_{n=1}^{\infty} \alpha_n(a) = \infty, \text{ and } \sum_{n=1}^{\infty} \alpha_n(a)^2 < \infty.$$
 (3)

The first condition ensures the steps are sufficiently large to overcome an initial condition and the second condition ensures that the steps become small enough to give convergence. However, step sizes which meet this conditions are rarely used in applications because they seem to converge slowly or require considerable tuning.

With this techniques, the initialisation effectively becomes an additional parameter which needs to be chosen. They can be used to provide prior knowledge and can encourage exploration, for example, by being optimistic. However, this sort of exploration is not useful for non-stationary problems (the task changes creates a renewed need for exploration).

The intuition behind UCB is that its better, when exploring, to select among the non-greedy actions which have the *potential* for being optimal. To do this, we maximise an estimate of the arm's value summed with some sort of uncertainty estimate. UCB often performs well but can be difficult to extent to RL from bandits.

We don't have to use value based methods. Instead, we could directly learn a preference over the actions and modify our preferences to give us higher rewards e.g., increasing our preference for an action if selecting it lead to a better outcome than expected.

Definition 2.2 (Contextual Bandits) In a contextual bandit, the agent is faced with a series of multi-armed bandit problems, each associated with some information i.e., the agent has information about which bandit problem is being faced at any timestep.

Contextual bandits are intermediate between multi-armed bandits and the full RL problem; they involve learning a policy (i.e., a mapping from state information to actions) but similar to k-armed bandits, **the action selected influences only the immediate reward and not the next situation**. If we drop this constraint, we have the full RL problem. UCB

Contextual Bandits

3 Finite Markov Decision Processes

Definition 3.1 (Finite Markov Decision Process (MDP)) A finite Markov Decision Process (MDP) is made up of:

- A sequence of discrete time steps, t = 0, 1, 2, ...
- A discrete set of states, $S_t \in S$.
- A set of actions which the agent can take, $A_t \in \mathcal{A}(S_t)$.
- A finite set of numerical rewards, $R_t \in \mathcal{R} \subset \mathbb{R}$. In this case, there is a well defined discrete transition distribution:

$$p(s', r|s, a) \triangleq Pr[S_t = s', R_t = r|S_{t-1} = s, A_{t-1} = a],$$
 (4)

which defines the dynamics of the MDP.

Note the Markov assumption; probabilities over the next state and reward depend only on the current state and action. This can be interpreted as a restriction on the state - the state must include information about all aspects of the past agent-environment interaction which will affect the future.

Given p, we can compute the following functions of interest:

$$p(s'|s,a) \triangleq \Pr[S_t = s'|S_{t-1} = s, A_{t-1} = a],$$
 (5)

$$r(s,a) \triangleq \mathbb{E}[R_t | S_{t-1} = s, A_{t-1} = a], \tag{6}$$

MDP

Interpreting the Markov Assumption

and

$$r(s, a, s') \triangleq \mathbb{E}[R_t | S_{t-1} = s, A_{t-1} = a, S_t = s'].$$
 (7)

A general rule of thumb is that anything that cannot be changed *arbitarily* by the agent is considered to be outside of it and thus part of the environment. The agent-environment boundary represents the limit of the agent's absolute control, not of its knowledge.

The goal of the agent is formalised in terms of the reward signal which passes from the environment to the agent. This is one of the most distinctive features of RL.

All of what we mean by goals and purposes can be well though of as the maximisation of the expected value of the cumulative sum of a received scalar signal.

The reward signal is not the place to impart to the agent prior knowlege about *how* to achieve a task; if we do this, the agent may only achieve subgoals without actually acheiving the real goal. A better place to impart this knowledge is the initial policy or value function. How does this link to reward shaping?

Episodic tasks terminate at some timestep, *T*, whilst continuing tasks do not. In the first case, we simply use the (expected) sum of rewards whilst we introduce discounting in the infinite horizon case i.e.,

$$G_t \triangleq \sum_{k=0}^{\infty} \gamma^k R_{t+k+1}. \tag{8}$$

 G_t is the *discounted return*. The closer the value of γ is to 1, the more farsighted the agent is.

We convert episodic tasks to continuing tasks by introducting a special *absorbing state* which transitions only to itself and returning zero reward. We can write:

$$G_t \triangleq \sum_{k=t+1}^{T} \gamma^{k-t-1} R_k, \tag{9}$$

allowing for $T = \infty$ and $\gamma = 1$. The book says not both - but I think that should be allowed for an episodic task with absorbing state.

Definition 3.2 (Policy) A policy is a mapping from states to probabilities of selecting each possible action. $\pi(a|s)$ is the distribution over actions, A_t , given the current state, S_t .

Definition 3.3 (Value Function) The value function of state s under policy π is the expected return when starting in s and following π .

$$\nu_{\pi}(s) \triangleq \mathbb{E}_{\pi}[G_t | S_t = s] \tag{10}$$

Definition 3.4 (Action-Value Function) The action-value function of action a, state s under policy π is the expected return when starting in s and following π after first performing a.

$$q_{\pi}(s,a) \triangleq \mathbb{E}_{\pi}[G_t|S_t = s, A_t = a] \tag{11}$$

Value functions satisfy recursive relationships.

A policy, π , is better than policy π' iff $\nu_{\pi}(s) \ge \nu_{\pi'}(s) \forall s \in \mathcal{S}$ with a strict inequality for at least one state. Optimal policies are better than or equal to all other policies. There may be multiple optimal policies, but they all share the same state-value function,

$$\nu_*(s) = \max_{\pi} \nu_{\pi}(s), \text{ for all } s \in \mathcal{S}.$$
 (12)

The optimal action-value function is define analogously.

The Bellman optimality equation for the value function is:

$$\nu_*(s) = \max_{a \in \mathcal{A}(s)} \sum_{s' \in S, r \in \mathcal{R}} p(s', r|s, a)[r + \gamma \nu_*(s')], \tag{13}$$

which expresses the intuition that the optimal value of a state is equal to the expected return of the best action from that state. Similarly, for the action-value function,

$$q_*(s,a) = \sum_{s',r} p(s',r|s,a) [r + \gamma \max_{a' \in \mathcal{A}(s')} q_*(s',a')].$$
 (14)

Agent vs Environment

Reward Hypothesis

Discount Rates

Unifying Notation

Policy

Value Function

Action-Value Function

Optimal Policies

Bellman Optimality Equality

For a finite MDP, the Bellman optimality equations give a system of equations which can be solved for $v_*(s)$. Once we have the optimal value functions, it is easy to determine an optimal policy by acting greedily (or performing a one-step search). If you use v_* to select actions in the short-term, the greedy policy is optimal in the long term since this value function takes into account reward consequences. The optimal expected long-term return is locally and immediately available for each state so a one-step lookahead search yields optimal actions.

Note that the action-value function effectively caches the results of all of the one-step lookahead searches. This function can be maximised (with respect to the action) to find the optimal policy.

This all sounds great. However, we run into the following problems:

- We need to accurately know the dynamics of the environment.
- We need to have enough computational resources.
- The Markov property needs to hold.

4 Dynamic Programming

Dynamic Programming (DP) algorithms can be used to compute optimal policies given perfect models of the environment. However, they are important only theoretically because they are limited by imperfect models and finite compute.

Policy evaluation computes the state-value function for policy π . We turn the Bellman equation into an update rule:

$$\nu_{k+1}(s) \triangleq \sum_{a \in \mathcal{A}(s)} \pi(a|s) \sum_{s' \in \mathcal{S}, r \in \mathcal{R}} p(s', r|s, a)[r + \gamma \nu_k(s)], \quad \forall s \in \mathcal{S}.$$
 (15)

 v_{π} is a fixed point for this update rule since it must satisfy the Bellman equation. Note that the estimate in general converges as $k \to \infty$ provided $\gamma < 1$ or eventual termination from all states under π (the same conditions which give a unique value function).

This is an *expected update*; the old value is replaced with a new value computed using previous values and expected immediate reward. In fact, all DP updates are called *expected updates* since they take the expectation over successor states rather than sampling.

Algorithm 1 Iterative Policy Evaluation

Input: π , the policy to be evaluated

Parameters: $\theta > 0$, a threshold determining accuracy.

Initialisation: initialise V(s) arbitarily for all $s \in S$ except that V(terminal) = 0

```
1: loop:

2: \Delta \leftarrow 0

3: for each s \in \mathcal{S} do: \triangleright Sweep over states, using the most recent values of V(s)

4: v \leftarrow V(s)

5: V(s) \leftarrow \sum_{a \in \mathcal{A}(s)} \pi(a|s) \sum_{s' \in \mathcal{S}, r \in \mathcal{R}} p(s', r|s, a)[r + \gamma V(s')]

6: \Delta \leftarrow \max(\Delta, |v - V(s)|)

7: until \Delta < \theta
```

If we know the dynamics, knowing the value of the policy helps us improve the policy. We can compute the value of selecting action a in state s and thereafter following π . If it is better to select a once in s and thereafter follow π , it is better to always select a in s and the new policy is an improvement.

Theorem 4.1 (Policy Improvement Theorem) *Let* π *and* π' *be any pair of determinstic policies such that, for all* $s \in S$,

$$q_{\pi}(s, \pi'(s)) \ge \nu_{\pi}(s). \tag{16}$$

Then π' must be as good as, or better than, π i.e., it must obtain greater or equal expected return from all states $s \in S$:

$$\nu_{\pi}'(s) \ge \nu_{\pi}(s). \tag{17}$$

Note that if there is any strict inequality in Eq. (16), there must be a strict inequality in Eq. (17) in at least one state.

Caveats

Limited Utility

Policy Evaluation

Policy Improvement Theorem

The greedy policy i.e., $\pi'(s) = \arg\max_a q_\pi(s,a)$, meets the conditions of the policy improvement theorem. The process of making a new policy which improves on an original policy by making it greedy with respect its value function is known as *policy improvement*. If policy improvement gives us a policy with the same perform as the old policy, this policy must necessarily be optimal as only the optimal policy satisfies the Bellman optimality equation.

Policy Improvement

Note that similar analysis can be performed for stochastic policies.

Policy Iteration

We can repeatedly evaluate policy π to calculate ν_{π} and then improve π to form π' . This process is guaranteed to find the optimal policy.

Algorithm 2 Policy Iteration

Initialisation: initialise V(s) arbitarily for all $s \in S$ except that V(terminal) = 0. Initalise $\pi(s) \in A(s)$ arbitrarily for all $s \in S$.

Parameters: $\theta > 0$, a threshold determining accuracy.

```
▶ Policy Evaluation
 1: loop:
             \Delta \leftarrow 0
 2:
            for each s \in \mathcal{S} do:
 3:
                  \begin{array}{l} v \leftarrow V(s) \\ V(s) \leftarrow \sum_{a \in \mathcal{A}(s)} \pi(a|s) \sum_{s' \in \mathcal{S}, r \in \mathcal{R}} p(s', r|s, a) [r + \gamma V(s')] \\ \Delta \leftarrow \max(\Delta, |v - V(s)|) \end{array}
  4:
  5.
 7: until \Delta < \theta
                                                                                                                    ▶ Policy Improvement
 8: policy-stable ← True
 9: for each s \in \mathcal{S} do:
            old-action \leftarrow \pi(s)
10:
             \pi(s) \leftarrow \arg\max_{a \in \mathcal{A}(s)} \sum_{s' \in \mathcal{S}, r \in \mathcal{R}} p(s', r|s, a) [r + \gamma V(s')]
11:
            if old-action \neq \pi(s) then:
12:
                  policy-stable ← True
13:
14: if policy-stable then:
            return V \simeq \nu_*, \pi \simeq \pi *
15:
16: else:
17:
              go to 1
```

It turns out that we don't even need to perform policy evaluation until convergence. We form the *value iteration* algorithm by performing one step of policy evaluation followed by policy improvement.

Value Iteration

Algorithm 3 Value Iteration

Initialisation: initialise V(s) arbitrarily for all $s \in \mathcal{S}$ except that V(terminal) = 0. Initialise $\pi(s) \in \mathcal{A}(s)$ arbitrarily for all $s \in \mathcal{S}$.

Parameters: $\theta > 0$, a threshold determining accuracy.

```
1: loop:

2: \Delta \leftarrow 0

3: for each s \in S do:

4: v \leftarrow V(s)

5: V(s) \leftarrow \max_{a \in A(s)} \sum_{s' \in S, r \in \mathcal{R}} p(s', r|s, a)[r + \gamma V(s')]

6: \Delta \leftarrow \max(\Delta, |v - V(s)|)

7: until \Delta < \theta

8: return \pi(s) = \arg\max_{a} p(s', r|s, a)[r + \gamma V(s')] \simeq \pi^*(s), V(s) \simeq \nu_*(s)
```

The algorithm can also be interpreted as the conversation of the Bellman optimality equality to an update rule. Above, V will converge to v_* .

Asynchrononous DP algorithms are in-place iterative DP algorithms which are not organised in terms of sweeping the entire state space (a drawback of normal DP). The algorithm *must* continue to update all states to ensure correct convergence.

Generalied policy iteration (GPI)¹ refers to the general idea of letting policy-evaluation

Alternative Interpretation

Asynchronous DP

Generalised Policy Iteration

and policy-improvement processes interact, independent of the granularity. If **both the evaluation and improvement stabilise, the value function and policy must be optimal**. The value function stabilises only when it is consisent with the current policy and policy stabilises only when it is greedy with respect to the current value function. Both stabilising implies that a policy has been found which is greedy with respect to its own evaluation function, implying that the Bellman optimality equation holds.

Whilst not practical for large problems, DP methods are actually quite efficient with (worst-case) complexity $\Omega(\text{poly}(n,k))$ where n,k are the number of states and actions respectively. They also seem to typically converge must faster than the worst case.

DP methods update estimates of the values of states based on estimates of the values of successor states; **bootstrapping**.

v,π v_*,π_* v_*,π_*

Generalised Policy Iteration (GPI)

Bootstrapping

5 Monte Carlo Methods

We now **relax the assumption that we have complete knowledge of the environment**, transforming the task into *learning* rather than *planning*. Whilst in DP, value functions were computed, here they will be learnt. Additionally, note that surprisingly often, even when we have perfect knowledge of the environment, it is hard to apply DP since the dynamics function, p, is difficult to obtain (though it can be easy to sample from).

Monte Carlo (MC) methods solve RL problems by averaging sample returns. Here, they are defined only for episodic tasks to ensure the returns are well defined. They are incremental on an episode-by-episode sense but not in a step-by-step sense.

*Monte Carlo Prediction*¹ attempts to learn the state-value function for a policy. An obvious way to estimate it is to simply average the returns observed after visiting that state.

Algorithm 4 First-visit MC Prediction

Input: policy π to be evaluated.

Initialisation: initialise V(s) arbitarily for all $s \in S$ except that V(terminal) = 0. Initialise returns $[s] \leftarrow []$ for all $s \in S$

```
1: loop:
        Generate an episode under \pi: \{S_0, A_0, R_1, S_1, A_1, R_2, ..., S_{T-1}, A_{T-1}, R_T\}
2:
3:
        for t \in \{T-1, T-2, ..., 0\} do:
4:
            G \leftarrow R_{t+1} + \gamma G
                                                                                 ▶ Recusively compute G
5:
            if S_t \notin \{S_0, S_1, \dots S_{t-1}\} then:
                                                                                               ▶ If first visit
6:
                append G to returns[S_t]
7:
                V(S_{t+1}) \leftarrow average(returns[S_t])
8:
```

The *every-visit* MC algorithm instead averages the returns following all visits to each state. Both every-visit and first-visit converge to ν_{π} quadratically as the number of visits to each state goes to infinity.

MC methods do not bootstrap; the estimates for each state are independent. Note that the computational expense of estimating the state value is independent of the total number of states and we can generate many episodes starting from states of interest if we desire.

Note that since we no longer have a model, knowing ν_{π} is not sufficient for policy improvement. We instead need to calculate q_{π} . This can be done in a manner similar to Algorithm 4 except that now each 'visit' is a tuple (s,a). However, there is no guarantee that each state-action pair is visited i.e., we have an issue of *maintaining exploration*. One way to resolve this is *exploring starts* where it is specified that the episode begins in a specific state-action pair and that every pair has a non-zero probability of selection. Alternatively, we could consider policies which always assign strictly positive probabilities to all actions.

We can naturally apply GPI, using MC methods to calculate q_{π} .

Compared to DP

Monte Carlo Methods

Monte Carlo Prediction



Monte Carlo Backup Diagram

Exploring Starts

Monte Carlo Control

¹Margin figure reproduced from Sutton, Richard S., and Andrew G. Barto. *Reinforcement learning: An introduction*. MIT press, 2018.

Algorithm 5 MCES: Monte Carlo Exploring Starts

Initialisation: initialise $\pi(s) \in \mathcal{A}(s)$ arbitarily, for all $s \in \mathcal{S}$. Initialise Q(s, a) arbitarily for all $s \in \mathcal{S}$, $a \in \mathcal{A}$. Initalise returns $[s, a] \leftarrow []$ for all $s \in \mathcal{S}$

```
1: loop:
          Choose S_0 \in \mathcal{S}, A_0 \in \mathcal{A}(S_0) such that all pairs have probability > 0
 2:
          Generate an episode from S_0, A_0 under \pi: \{S_0, A_0, R_1, S_1, A_1, R_2, ..., S_{T-1}, A_{T-1}, R_T\}
 3:
 4:
          for t \in \{T-1, T-2, ..., 0\} do:
 5:
              G \leftarrow R_{t+1} + \gamma G
                                                                                        ▶ Recusively compute G
 6:
              if (S_t, A_t) \notin \{(S_0, A_0), (S_1, A_0), \dots (S_{t-1}, A_{t-1})\} then:
                                                                                                       ▶ If first visit
 7:
                   append G to returns[S_t, A_t]
 8:
                   Q(S_{t+1}, A_t) \leftarrow average(returns[S_t])
 9:
                   \pi(S_t) \leftarrow \arg\max_{a \in \mathcal{A}(S_t)} Q(S_t, a)
10:
```

Note that all returns for each state-action pair are averaged regardless of the policy used. MCES cannot converge to any suboptimal optimal; if it did, the value function would eventually converge to the correct value function, which would cause the policy to change if it is suboptimal. Additionally, it seems that updates to Q ought to decrease over time. However, formal correct convergence has not been proved for this algorithm.

Exploring starts cannot be relied upon in general, especially when learning directly from actual interaction with an environment. We need to relax this, which we can do by, in general, allowing the agent to continue selecting them.

Relaxing Exploring Starts

Algorithm 6 On-policy first-visit MC Control

```
Parameter:
```

small $\epsilon > 0$

Initialisation:

```
\pi as an arbitarily \epsilon-soft policy.
Q(s, a) arbitarily for all s \in \mathcal{S}, a \in \mathcal{A}.
returns [s,a] \leftarrow [] for all s \in S
```

```
1: loop:
```

```
2:
            Generate an episode from under \pi: \{S_0, A_0, R_1, S_1, A_1, R_2, ..., S_{T-1}, A_{T-1}, R_T\}
 3:
            for t \in \{T-1, T-2, ..., 0\} do:
 4:
                  G \leftarrow R_{t+1} + \gamma G
                                                                                                              ▶ Recusively compute G
 5:
                  if (S_t, A_t) \notin \{(S_0, A_0), (S_1, A_0), \dots (S_{t-1}, A_{t-1})\} then:
                                                                                                                                  ▶ If first visit
 6:
                        append G to returns [S_t, A_t]
 7:
                        Q(S_t, A_t) \leftarrow average(returns[S_t, A_t])
 8:
                       A^* \leftarrow \arg\max_{a \in \mathcal{A}(S_t)} Q(S_t, a)
\pi(a|S_t) \leftarrow \begin{cases} 1 - \epsilon + \epsilon/\mathcal{A}(S_t), & a = A^* \\ \epsilon/\mathcal{A}(S_t), & \text{otherwise} \end{cases}
                                                                                                                  ▶ Break ties arbitarily
 9:
                                                                                                          \forall a \in \mathcal{A}(S_t)
10:
```

The policy improvement theorem ensures that any ϵ -greedy policy is an improvement over any ϵ -soft policy. Policy iteration works for ϵ -soft policies, and the above algorithm will find the best possible soft policy.

Learning control methods aim to learn action values conditional on subsequent optimal behaviour but they need to behave non-optimally to explore all actions. Algorithm 6 is a compromise, learning action-values for a 'near'-optimal policy. Alternatively, we could use two policies; one to generate behaviour (the behavioural policy) and one to approach optimal behaviour (the target policy). This approach is known as off-policy learning.

The *prediction* policy seeks to estimate q_{π} or v_{π} but from behaviour generated by policy b. We need to make the assumption of **coverage**, $\pi(a|s) > 0 \Rightarrow b(a|s) > 0$. We do offpolicy learning most frequently using importance sampling (IS). Sparing the derivation, the ordinary importance sampling estimator is:

Off-Policy Learning

Ordinary Importance Sampling

$$\nu_{\pi}(s) = \mathbb{E}_{\pi}[G_t | S_t = s]$$

$$\simeq \sum_{t \in \mathcal{T}(s)} \frac{\rho_{t:T(t)-1}}{|\mathcal{T}(s)|} G_t$$
(18)

where, for notational convenience, the episodes are appended to one another so that a timestep uniquely indentifies a timestep within an episode, $\mathcal{T}(s)$ is the set of times at which s was visited, T(t) is the termination time of the episode which t belong to. $\rho_{t,T-1}$ are the importance weights:

$$\rho_{t:T-1} = \prod_{k=t}^{T-1} \frac{p(R_{k+1}, S_{k+1}|S_k, A_k)\pi(A_k|S_k)}{p(R_{k+1}, S_{k+1}|S_k, A_k)b(A_k|S_k)}$$

$$= \prod_{k=t}^{T-1} \frac{\pi(A_k|S_k)}{b(A_k|S_k)}.$$
(19)

The above formulae are straightforward to derive following the standard importance sampling derivation. Note that the dynamics of the unknown MDP cancel out.

An alternative to Eq. (20) is weighted importance sampling:

$$v_{\pi}(s) \simeq \sum_{t \in \mathcal{T}(s)} \frac{\rho_{t:T(t)-1}}{\sum_{t \in \mathcal{T}(s)} \rho_{t:T(t)-1}} G_t.$$
 (20)

The first-visit version of the ordinary IS estimator is unbiased with high variance (since the variance of the importance weights can be unbounded) whilst the first-visit version of the weighted IS estimator is biased but with much lower variance: the maximum weight on any of the returns is 1. In fact, assuming bounded returns, the variance of the weighted IS estimator converges to zero even if the importance weights have infinite variance. In practice, the weighted estimator has much lower variance and is prefered. Additionally, note that the weighted importance sampling estimator produces a weighted average of only the returns consisent with the target policy whilst the normal importance sampling estimate does not.

The every-visit methods introduce bias, but the bias asymtoticallty falls to zero as the number of samples increases. These methods are prefered as they remove the need to track which states have been visited and they are easier to extend.

First vs. Every Visit

Weighted Important Sampling

Algorithm 7 Off-policy Every-Visit MC Prediction

Input:

arbitary target policy π

Initialisation:

```
Q(s, a) arbitarily for all s \in \mathcal{S}, a \in \mathcal{A}.
C(s,a) \leftarrow 0 for all s \in \mathcal{S}, a \in \mathcal{A}
```

```
1: loop:
```

```
2:
             b \leftarrow any policy which coverage of \pi.
             Generate an episode from under b: \{S_0, A_0, R_1, S_1, A_1, R_2, ..., S_{T-1}, A_{T-1}, R_T\}
 3:
 4:
             W \leftarrow 1
  5:
             for t \in \{T-1, T-2, ..., 0\} do:
  6:
                  G \leftarrow R_{t+1} + \gamma G
C(S_t, A_t) \leftarrow C(S_t, A_t) + W
Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \frac{W}{C(S_t, A_t)} [G - Q(S_t, A_t)]
W \leftarrow W \frac{\pi(A_t | S_t)}{b(A_t | S_t)}
if W = 0 then:
                                                                                                                       ▶ Recusively compute G
  7:
 8:
 9:
10:
11:
                          break
12:
```

Algorithm 8 Off-policy MC Control

```
Initialisation:
```

```
Q(s, a) arbitarily for all s \in \mathcal{S}, a \in \mathcal{A}.
              C(s,a) \leftarrow 0 for all s \in \mathcal{S}, a \in \mathcal{A}
              \pi(s) \leftarrow \arg \max_a Q(s, a), with ties broken consistently.
 1: loop:
 2:
             b \leftarrow \text{any soft policy}
             Generate an episode from under b: \{S_0, A_0, R_1, S_1, A_1, R_2, ..., S_{T-1}, A_{T-1}, R_T\}
 3:
  4:
             W \leftarrow 1
 5:
             for t \in \{T-1, T-2, ..., 0\} do:
 6:
                   G \leftarrow R_{t+1} + \gamma G
                                                                                                                     ▶ Recusively compute G
 7:
                   \begin{split} &C(S_t, A_t) \leftarrow C(S_t, A_t) + W \\ &Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \frac{W}{C(S_t, A_t)} [G - Q(S_t, A_t)] \\ &\pi(S_t) \leftarrow \arg\max_{a} Q(S_t, a) \text{ with ties broken consistently.} \end{split}
 8:
 9:
10:
                   if A_t \neq \pi(S_t) then:
11:
                         break
12:
                   else
13:
                         W \leftarrow \frac{W}{b(A_t|S_t)}
14:
```

Algorithm 8 shows an approach to off-policy MC control using weighted importance sampling.

A problem with this algorithm is that it learns only from the tails of episodes where the remaining actions are greedy with respect to the current policy. If non-greedy actions are common, learning will be slow, especially for early actions. Off-Policy MC Control

6 Temporal-Difference Learning

Temporal-Difference Learning (TD Learning) is a combination of Monte Carlo and Dynamic Programming ideas. Like MC methods, TD learning uses raw experience without an environmental model but it also bootstraps, using estimates of successor states to improve state estimates.

Monte Carlo methods wait until the return following the visit is known and then use return as a target for $V(S_t)$. For example, *constant-* α *MC* uses:

$$V(S_t) \leftarrow V(S_t) + \alpha [G_t - V(S_t)], \tag{21}$$

which is simple and appropriate for non-stationary problems. TD methods wait only until the next time step, bootstrapping to estimate the reward,

$$V(S_t) \leftarrow V(S_t) + \alpha [R_{t+1} + \gamma V(S_{t+1}) - V(S_t)]. \tag{22}$$

This is known as the TD(0) update, an estimate of the value of state as it samples the reward and uses an estimate of the successor state.

Temporal-Difference Learning

TD Prediction



TD(0) Backup Diagram

Algorithm 9 Tabular TD(0) Prediction

Input:

policy π to be evaluated. step size $\alpha \in [0, 1]$

Initialisation:

initialise V(s) arbitarily for all $s \in \mathcal{S}$ except that V(terminal) = 0. Initalise

- 1: **for** each episode **do**:
- 2: Generate initial state *S*.
- 3: **while** episode not terminated **do**:
- 4: $A \leftarrow \pi(S)$
- 5: Perform action A, observe R, S'
- 6: $V(S) \leftarrow V(S) + \alpha [R + \gamma V(S') V(S)]$
- 7: $S \leftarrow S$

The TD error is defined as:

TD Error

$$\delta_t \triangleq R_{t+1} + \gamma V(S_{t+1}) - \gamma V(S). \tag{23}$$

The RD error is the error in the estimate *made at that time*. Note that δ_t is only available at time t+1. It can be shown that the Monte Carlo error can be written as a sum of TD errors:

$$G_t - V(S_t) = \sum_{k=t}^{T-1} \gamma^{k-t} \delta_k$$
 (24)

TD methods can be implemented in an online, fully incremental fashion. This is beneficial because many tasks have very long episodes which result in MC methods learning slowly. MC methods often have to neglect returns gained from actions which are inconsisent with the policy we are attemping to evaluate, but since TD learning uses *transitions*, it is much less suspectible to this problem.

Note that it can be shown that TD(0), given a reducing step-size parameter satisfying the typical stochastic optimisation conditions, converges with probability one in the mean. Even though bootstrapping might not seem sound, it seems to work. Additionally, these methods often converge faster than MC methods.

Batch updating is when the update is computed across a batch of training episodes, the value function only being updated once for each batch. For small enough α , batch α -MC converges to the sample averages of actual returns.

Advantages of TD Prediction

Batch Updating

Temporal-Difference vs Monte Carlo Prediction

We are faced with a *prediction problem* in an unknown Markov reward process without discounting. Suppose that we observe the following episodes.

 A, 0, B, 0
 B, 0
 B, 1
 B, 1

 B, 1
 B, 1
 B, 1
 B, 1

where, for example, the first episode indicates that we started in state A, transitioned to state B with reward 0 etc. It is clear that the value of state B is 0.75, as this is the frequency that B returned reward 1 before termination. What about state A?

TD(0) would estimate of V(A) as 0.75; it would observe that every state the MRP was in state A, it transitioned to B and the value of B is 0.75. This can be understood as using the MRP assumption. This does not minimise error on the training episodes, but we might expect this to be produce lower error on *future error*. In fact, this prediction is the value for the *maximum-likelihood model of the MRP*, also known the *certainty-equivalence estimate*, as it assumes that this model is the true model.

MC would instead note that every time the MRP was in state A, the return was 0 and estimate V(A) as 0. Note that this answer gives minimum squared error on the training error.

The non-batch version of these algorithms can be understood to be moving in these general directions, whilst not getting fully there. Note that with state space $n = |\mathcal{S}|$, the maximum-likelihood MRP requires up to n^2 memory to compute the transition probabilities, and then computing the value function requires an additional order n^3 steps (if done conventionally). It is striking that TD methods can approximate this solution in order n.

To solve the control problem, we must estimate action-values. Instead of considering transitions from states to states, we consider transitions from state-action pairs to state-action pairs, which again gives a Markov chain with a reward process.

$$Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \alpha [R_{t+1} + \gamma Q(S_{t+1}, A_{t+1}) - Q(S_t, A_t)]$$
 (25)

Sarsa

Algorithm 10 On-policy TD Control

Parameters:

```
step size \alpha \in (0,1]
small \epsilon > 0
```

Initialisation:

Q(s,a) arbitarily for all $s \in \mathcal{S}, a \in \mathcal{A}$ except that $Q(\texttt{terminal}, \cdot) = 0$

```
1: loop for each episode:
```

```
2: initialise S
```

3: Choose *A* using $Q(S, \cdot)$ e.g., ϵ -greedy.

4: **loop** for each episode step:

5: Take action A, observe S', R

6: Choose A' using $Q(S', \cdot)$

7: $Q(S,A) \leftarrow Q(S,A) + \alpha [R + \gamma Q(S',A') - Q(S,A)]$

8: $S \leftarrow S'$

9: $A \leftarrow A'$

Algorithm 10 shows a control algorithm using Sarsa. As in all on-policy methods, we continually estimate q_π for the behavioral policy π and update π towards greediness. This algorithm converges to the optimal policy and action-value function provided all state-action pairs are visited infinitely often and the policy converges to the greedy policy. *Q-Learning* is off-policy TD Control.

$$Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \alpha [R_{t+1} + \gamma \max_{a} Q(S_{t+1}, A) - Q(S_t, A_t)].$$
 (26)

The learned action-value function directly approximates q_* , while the behavioural policy determines which state-action pairs are visited. As usual, we require that all state-action pairs are visited for correct convergence.

Expected Sarsa instead uses the expected value over the successor state-action pairs:

$$Q(S_t, A_t) \leftarrow Q(S_t, A_t) + \alpha [R_{t+1} + \gamma \sum_{a} \pi(a|S_{t+1}) \ Q(S_{t+1}, a) - Q(S_t, A_t)]. \tag{27}$$

This algorithm moves deterministically in the same direction as Sarsa moves in expectation. Whilst more complex than Sarsa, it reduces the variance due to the noise in the selection of A_{t+1} , and given the same amount of experience, it tends to perform better than Sarsa. The reduction in variance often allows larger values of α to be used.

Note that the Expected Sarsa algorithm subsumes Q-learning; the target policy can be different from the behavioural policy.

The control algorithms discussed involve maximisation to construct target policies; a maximisation over expected values is used implicitly as an estimate of the maximum value, leading to significant positive bias. One way to view this problem is that this occurs because the same samples have been used for *both* determining the best action *and* estimating its value.

Double Learning attempts to reduce maximisation bias by instead maintaining two sets of value functions; one used to estimate the value, and the other used to estimate the optimal action. At each time-step, we update one set of values only. The values updated are often chosen with a coin-toss, and the behavioural policy could for example by the ϵ -greedy policy using the sum of the independent Q values.

Afterstates

In certain situations, initial parts of the environment dynamics are known, but not necessarily the full dynamics. This is common in games, such as chess or tic-tac-toe. In these situations, it is often beneficial to place value functions over *afterstates* i.e., the state after the known part of the dynamics function is applied. This reduces the size of the problem, as many state-action pairs have the same afterstates, and thus ought to have the same *afterstate value function*.

Generalised policy iteration can be applied as usual in this setting.

Q-Learning

Expected Sarsa

Maximisation Bias

Double Learning